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# **EUROPEAN PATENT APPLICATION**

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(54) Olefin polymerization catalyst and process for producing polyolefin using the same

(57) A catalyst system comprising a transition metal compound (A) having at least one of a cyclopentadienyl group or a substituted cyclopentadienyl group and at least one of a cyclic ligand containing a hetero atom and having a delocatised x bond, and an organoalluminum compound (B), or (A), (B) and a compound (C) forming an ionic complex by reacting with a transition metal compound; and a process for preparing an oletin polymer using said catalyst system.

### Description

### BACKGROUND OF THE INVENTION

#### 5 Field of the Invention

The present invention relates to a catalyst to polymerizing an olefin and a process for preparing a pobylefin. In particular, the present invention relates to a catalyst having a superior copplymerability and a process for preparing a loyelefin having a proper molecular weight and a high content of comonomer, and particularly, a linear low density poltory weight and the proper molecular weight and a high content of comonomer, and particularly, a linear low density polter weight and the proper molecular weight and the proper prope

### Prior Art

Many reports have been published concerning a process for preparing a polyolefin with a metallocene catalyst. For example, in Japanese Patent Publication (Unexamined) No. Sho 59-19309(1983), a process for producing a polyolefin by using a metallocene catalyst and alturinoxane is disclosed. However, this method, namely, copolymetration of ethylene and an a-client performed with a system using bis(polpotestadieny) zirconium dichibride and methyl aluminoxane provides a polymer having a low modecular weight and a low content of a-cliefin. In Japanese Patent Publication (Nexamined) No. Hel 1-502056(1989), it is disclosed that ethylene-c-cliefin copolymetration by using a metallocene catalyst and a born compount, for example, bis(perhametryl originationy)dimetryl circonium and trifu-buyl)parmonium tetra[pentaltu-copheny) borate, is performed, but only a copolymer having a low molecular weight and a low content of ca-clief in is challend.

On the other hand, in Japanese Patent Publication (Unexamined) Hei No. 3-163098(1991), it is disclosed that ethylene-a-clefin copylemerization is performed by using (lert-butylamide)dimethyl(tetra methyl-n<sup>5</sup>-cyclopentadienyl) silane titanium dichloride, but as a result, a polymer having a higher molecular weight than an ethylene-a-clefin copylymer prepared by the metallocene catalyst as described above is obtained. However, this is still insufficient. In Japanese Patent Publication (Unexamined) No. Hei 6-49103(1994), it is disclosed that ethylene-a-clefin copylomerization is performed by using bis(2,3,4,5-tetramethylphosphoryl) zirconium dichloride but a polymer having a low molecular weight, a broad molecular weight distribution and a high melting point is still provided. As described above, eliviene-a-defin copylomers obtained by the use of a conventional metallocene catalyst have a low molecular weight.

### SUMMARY OF THE INVENTION

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It is an object of the present invention to provide a catalyst giving an olefin polymer having a high molecular weight and a process for producing a polydefin by using the same. The use of item "olefin polymer" herein indicates both a homopolymer of an olefin or a copolymer of an olefin with another olefin.

The present invention relates to a catalyst component comprising a compound (A) described as follows, a catalyst system comprising the compound (A) and a compound (B) or the compound (A), the compound (B) and a compound (C) as main components and a process for producing an olefin polymer by using these catalyst systems.

Compound(A): a transition metal compound represented by the general formula (1) having at least one of a cyclopentationly group or a substituted cyclopentationly group and at least one of a cyclic ligand containing a hetero atom and having a delocalized x bond,

$$\begin{pmatrix}
R^{1} & R^{1} \\
R^{1} & C & R^{1} \\
C & C & R^{1}
\end{pmatrix}
\begin{pmatrix}
R^{2}_{a} & X & R^{2}_{a} \\
R^{2}_{a} & X & X^{2}_{a}
\end{pmatrix}_{m} MY_{n} \quad (1)$$

wherein M represents an element of the Third Group, the Fourth Group or lanthanide series in the Periodic Table,

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each of R<sup>1</sup> and R<sup>2</sup> independently represents a hydrogen atom, a halogen atom, a hydrocarbon group for a halogenated hydrocarbon group having 1 to 20 carbon atoms, wherein R<sup>1</sup> and R<sup>2</sup> may be substituted with a group containing a nitrogen atom, a phosphorus atom, an oxygen atom, a suffur atom or a silicon atom, all of R<sup>2</sup> and R<sup>2</sup> may be the same or different, groups R<sup>1</sup> together with the carbon atoms to which they are attached, and generally adjacent to each other, may form a ring, generally a 5- or 6-membered ring, gones R<sup>2</sup> together with the groups X to which they are attached, and generally adjacent to each other, may form a ring, generally a 5- or 6-membered ring, and R<sup>2</sup> and R<sup>2</sup> may be bonded together; substituent X indicates an element of the 13th Corup, 14th Group or 15th Group in the Periodic Table, each substituent X may be the same or different but at least one substituent X is an element other than a carbon atom; Y is a hydrogen atom, a halogen atom or a hydrocarbon group having 1 to 20 carbon atoms; I, mand n are integers, 13tl63, 1sm3 and 05m2, and a 6 to 7 1; and when Mis a metal of the third group or a lantharide series element, I = m + m = 3 and when Mis a metal of the Fourth Group, I + m + m = 4. (8) an oranoallastimum compound as elected from the following commounds:

(B1): an organoaluminum compound represented by the general formula R3bAIZ3b

(B2): a cyclic aluminoxane (B2a) having a structure represented by the general formula {-Al(R<sup>4</sup>)-O-}<sub>2</sub> and/or a linear aluminoxane (B2b) having a structure represented by the general formula R<sup>5</sup>{-Al(R<sup>5</sup>)-O-}<sub>3</sub>AlR<sup>5</sup><sub>2</sub>

wherein  $R^2$ ,  $R^4$  and  $R^5$  represent a hydrocarbon group having 1 to 8 carbon atoms. In the above general formula, all of  $R^5$  and all of  $R^5$  may be the same or different. Z is a hydrogen and/or a halogen, b is a number of 0 to 3, and cand of represent an integer of 1 or more.

(C): a compound forming an ionic complex by reacting with a transition metal compound.

In this specification including the accompanying claims references to the groups of the Periodic Table are to be understood as follows: the Third Group comprises scandium and ythium; the Fourth Group comprises thanium, zirconium and hathium; the 13th Group comprises boron; the 14th group comprises carbon and silicon; and the 15th Group comprises nitroen and phosphorus.

The present invention is further explained in detail as follows.

Figure 1 at the end of the description, is a flow chart drawing to aid the comprehension of the present invention and shows one of the presentative embodiments of the present invention.

### DETAILED DESCRIPTION OF THE INVENTION

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- 1) The transition metal catalyst component for polymerizing an olefin.
- In the present invention, the compound (A) is the compound represented by the general formula (1).

$$\begin{pmatrix}
R^{1} & R^{2} & R^{2} \\
R^{1} & R^{1}
\end{pmatrix}
\begin{pmatrix}
R^{2} & X & R^{2} \\
R^{2} & X & X^{2} \\
R^{2} & X & R^{2} \\
R^{2} & X & R^{2}
\end{pmatrix}_{m} MY_{n} \quad (1)$$

wherein M represents an element of the Third Group, the Fourth Group or lamthaniste series in the Periodic Table, each of R<sup>1</sup> and R<sup>2</sup> independently represents a phytogen atom, a haldgein atom, a hydrocarbon group or a halogen atom hydrocarbon group having 1 to 20 carbon atoms, wherein R<sup>1</sup> or R<sup>2</sup> may be substituted with a group containing a nitrogen atom, a phosphorus atom, an oxygen atom, a suffur atom or a silicon atom, all of R<sup>1</sup> and R<sup>2</sup> may be the same or different, each of R<sup>1</sup> and seen of R<sup>2</sup> may be bonded together. X indicates an element of the 13th Group, 14th Group or 15th Group in the Periodic Table, each of X may be been same or different but at least one of the X's contains an element of the rath as a carbon atom. Y is a hydrogen atom, a halogen atom or a hydrocarbon group having 1 to 20 carbons. L m and n are integers, 1sts3, 1sm32 and 0sm32 and also or 1. When M is a metal of the Tritid Group or a lanthands essive selement. I = n n - 3 and when M is a metal of the Tritid Group or a lanthands essive selement. I = n n - 3 and when M is a metal

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of the Fourth Group, 1+ m + n + 4. Min the compound (A) indicates an element of the Third Group, the Profith Group, or the Lanthanide Series in the periodic table. The example of M includes a scandium or ythrium atom as a transition metal element of the Third Group in the Periodic Table; a titanium atom, a zirconium atom or a hafnium atom as a transition metal element of the Fourth Group; a samarium atom or the like as a transition metal element of the Lanthanide Series. A titanium atom. a zirconium atom or a thermium atom or a transition matom or a transition atom or a thermium atom or a transition.

Each of R<sup>1</sup> and R<sup>2</sup> in the compound (A) is a hydrogen atom, a halogen atom, a hydrocarbon group or a halogen atom, a hydrocarbon group having 1 to 2 clarbons, which may be substitlated with a group containing a integer atom, a phosphous atom, an oxygen atom, a suffur atom or a silicon atom, all of R<sup>1</sup> and R<sup>2</sup>may be the same or different, each of R<sup>1</sup> and each of R<sup>2</sup> may be bonded toolether to form a cucilic ring and R<sup>2</sup> and R<sup>2</sup> may be bonded toolether.

The example of R<sup>1</sup> and R<sup>2</sup> includes a fluorine atom, a chlorine atom, a bromine atom or an locline atom as a halogen atom, methyl group, ethyl group, n-propyl group, n-buyl group, n-buyl group, s-buyl group, s-buyl group, n-pentyl group, nepentyl group, n-beyl group, pnehyl group, bendyl group, bendyl group, pentaltuorophenyl group, pentaltuorophenyl group or the like as a hydrocarbon group or a halogenated hydrocarbon group having 1 to 20 carbons. A hydrogen atom, methyl group, ethyl group, t-buyl group or n-buyl group is preferred.

Substituted groups containing a nitrogen atom, a phosphorus atom, an oxygen atom, a sulfur atom or a silicon atom include -NP<sup>6</sup>, -NP<sup>6</sup>, -NP<sup>8</sup>, -PP<sup>8</sup>, -DO, -OR<sup>10</sup>, -S, -SP<sup>11</sup>, -SR<sup>10</sup><sub>2</sub>, -SR<sup>10</sup>, -SR<sup>10</sup>, -R<sup>10</sup>, -R

The example of the group containing a nitrogen atom includes methylamino group, ethylamino group, n-propylamino group, isopropylamino group, n-butylamino group, sobutylamino group, s-butylamino group, t-butylamino group, n-bexylamino group, n-cotylamino group, phenylamino group, tilino-methylamino group, pentalfuropolen nylamino group, di-s-butylamino group, di-b-butylamino group, di-n-propylamino group, di-n-butylamino group,

The example of the group containing a phosphous atom includes methyphosphino group, ethylphosphino group, n-putyphosphino group, activity group, n-butyphosphino group, powerphosphino group, n-butyphosphino group, n-delyphosphino group, phenylphosphino group, phenylphosphino group, pitrallurophenyl phosphino group, diethylphosphino group, group, diethylphosphino group, bisterflationenthylphosphino group, diethylphosphino group, bisterflationenthylphosphino group, bisterflationenthylphosphino group, diethylphosphino group, bisterf

The example of the group containing an oxygen atom includes methoxy group, ethoxy group, n-propoxy group, isopropoxy group, n-butoxy group, iso-butoxy group, s-butoxy group, n-hexoxy group, n-octoxy group, phenoxy group, bifurormethoxy group, pentatiproxphenoxy group or the like.

The example of the group containing a sulfur atom includes methylsulfide group, ethylsulfide group, h-nyspylsulfide group, inchylsulfide group, phenylsulfide group, the group of the like.

The example of the group containing a silicon atom includes dimethylsilylene group, diethylsilylene group, di-n-proylsilylene group, discoproylsilylene group, di-n-day/silylene group, di-n-bay/silylene group, di-n-ba

Examples of the ligand wherein each of R1 groups and each of R2 groups are bonded to form a cyclic ring in the compound (A) include indenyl group, t-butyfindenyl group, methyltidenyl group, dimethyltidenyl group, tetrahydroindenyl group, methyltidenyl group, tetrahydroindenyl group, indenyl group, carbazolyl group, butyfilmodelyl group, tetrahydroindolyl group, group, methyldroindolyl group, carbazolyl group, butyfilmodelyl group, methyldrospholindenyl group, phospholindenyl group, phospholindenyl group, phospholindenyl group, phospholindenyl group, fluorenyl group, indenyl group, phospholindenyl group, phospholind

When R<sup>1</sup> and R<sup>2</sup> are bonded together, they can form a grouping - (CRI<sup>4</sup><sub>2</sub>)<sub>x</sub> - (SIRI<sup>5</sup><sub>2</sub>)<sub>y</sub> or the like. RI<sup>4</sup> and RI<sup>5</sup> are a 5 hydrogen, a saturated hydrocarbon, anyl and allyl group, having up to 20 carbons, and these groups are unsubstituted or substituted, and x and year integers of 1 or more.

The example of these groups includes methylene group, dimethylene group, diphenylmethylene group, ethylene group, tetramethylethylene group, tetraphenylethylene group, silylene group, dimethylsilylene group, diphenylsilylene group, disilylene group, tetramethyldisilylene group, tetraphenyldisilylene group or the like. A armethylsilylene group ethylene group or methylene group is preferred.

X in the compound (A) indicates an element of the 13th Group, 14th Group or 15th Group in the Periodic Table, each of X may be the same of different and at least one of the few 5x contains an element other than a cation atom. A born atom, a nitrogen atom or a phosphorus atom is preferred and among them a nitrogen atom, or a phosphorus atom is more preferred.

V in the compound (A) is a hydrogen atom, a halogen atom or a hydrocarbon group having 1 to 20 carbons. The example of Y includes a fluorine atom, a chlorine atom, a bromine atom or an iodine atom as a halogen atom, and a methyl group, ethyl group, n-propyl group, isopropyl group, n-butly group, phenyl group, betwy group or the like as a hydrocarbon group having 1 to 20 carbons. Y is preferably a halogen atom or alikyl group having 1 to 8 carbons. When he preferably Y is a chlorine atom. I, m and in in the compound (A) are integers and 1453.3, IsmiS3 and 05x162. When M is a metal of the Third Group or a Lanthanide Series element, I + m + n = 3 and when M is a metal of the Fourth Group, I + m + n = 4. a is 0 or 1.1 it is preferred that in is 2.

Examples of the above-mentioned compound represented as the compound (A) include cyclopentadienyl pyrrolyl titanium dichloride, cyclopentadienyl methylpyrrolyl titanium dichloride, cyclopentadienyl dimethylpyrrolyl titanium dichloride, cyclopentadienyl tetramethylpyrrolyl titanium dichloride, cyclopentadienyl t-butylpyrrolyl titanium dichloride, cyclopentadienyl di-t-butytpyrrolyl titanium dichloride, pentamethylcyclopentadienyl pyrrolyl titanium dichloride, pentamethylcyclopentadienyl methylpyrrolyl titanium dichloride, pentamethylcyclopentadienyl dimethylpyrrolyl titanium dichloride, pentamethylcyclopentadienyl tetramethylpyrrolyl titanium dichloride, pentamethylcyclopentadienyl t-butylpyrrolyl titanium dichloride, pentamethyloyolopentadienyl di-t-butyloyrrolyl titanium dichloride, indenyl pyrrolyl titanium dichloride. ride, indenvi methylpyrrolyl titanium dichloride, indenvi dimethylpyrrolyl titanium dichloride, indenvi tetramethylpyrrolyl titanium dichloride, indenyl t-butylpyrrolyl titanium dichloride, indenyl di-t-butylpyrrolyl titanium dichloride. cyclopentadienyl indolyl titanium dichloride, cyclopentadienyl methylindolyl titanium dichloride, cyclopentadienyl dimethylindolyl titanium dichloride, cyclopentadienyl t-butylindolyl titanium dichloride, cyclopentadienyl di-t-butylindolyl titanium dichloride, pentamethylcyclopentadienyl indolyl titanium dichloride, pentamethylcyclopentadienyl methylindolyl titanium dichloride, pentamethylcyclopentadienyl dimethylindolyl titanium dichloride, pentamethylcyclopentadienyl t-butylindolyl titanium dichloride, pentamethyloyolopentadienyl di-t-butylindolyl titanium dichloride, indenyl indolyl titanium dichloride, indenyl methylindolyl titanium dichloride, indenyl dimethylindolyl titanium dichloride, indenyl t-butylindolyl titanium dichloride, indenyl di-t-butylindolyl titanium dichloride, cyclopentadienyl tetramethyl phospholyl titanium dichloride, cyclopentadienyl tetraphenyl phospholyl titanium dichloride, cyclopentadienyl phosphoindenyl titanium dichloride, pentamethylcyclopentadienyl tetramethyl phospholyl titanium dichloride, pentamethylcyclopentadienyl tetraphenyl phospholyl titanium dichloride, pentamethylcyclopentadienyl phosphoindenyl titanium dichloride, indenyl tetramethyl phospholyl titanium dichloride, indenyl tetraphenyl phospholyl titanium dichloride, indenyl phosphoindenyl titanium dichloride and the like.

Such Examples also include methylenecyclopentadienyl pyrrolyl titanium dichloride, methylenecyclopentadienyl methylpyrrolyl titanium dichloride, methylenecyclopentadienyl dimethylpyrrolyl titanium dichloride, methylenecyclopentadienyl trimethylpyrrolyl titanium dichloride, methylenecyclopentadienyl t-butylpyrrolyl titanium dichloride, methylenecyclopentadienyl di-t-butylpyrrolyl titanium dichloride, methylenetetramethylcyclopentadienyl pyrrolyl titanium dichloride, methylenetetramethylcyclopentadienyl methylpyrrolyl titanium dichloride, methylenetetramethylcyclopentadienyl dimethylpyrrolyl titanium dichloride, methylenetetramethylcyclopentadienyl trimethylpyrrolyl titanium dichloride, methylenetetramethylcyclopentadienyl t-butylpyrrolyl titanium dichloride, methylenetetramethylcyclopentadienyl di-tbutylpyrrolyl titanium dichloride, methyleneindenyl pyrrolyl titanium dichloride, methyleneindenyl methylpyrrolyl titanium dichloride, methyleneindenyl dimethylpyrrolyl titanium dichloride, methyleneindenyl trimethylpyrrolyl titanium dichloride, methyleneindenyl t-butylpyrrolyl titanium dichloride, methyleneindenyl di-t-butylpyrrolyl di-t-butylpyrrolyl di-t-butylpyrrolyl di-t-butylpyrrolyl di-t-butylpyrrolyl di-t-butylpyrrolyl di-t-butylpyrrolyl di-t-butylpyrrolyl cyclopentadienyl indolyl titanium dichloride, methylenecyclopentadienyl methylindolyl titanium dichloride, methylenecy-45 clopentadienyl dimethylindolyl titanium dichloride, methylenecyclopentadienyl t-butylindolyl titanium dichloride, methylenecyclopentadienyl di-t-butylindolyl titanium dichloride, methylenetetramethylcyclopentadienyl indolyl titanium dichloride, methylenetetramethylcyclopentadienyl methylindolyl titanium dichloride, methylenetetramethylcyclopentadienyl dimethylindolyl titanium dichloride, methylenetetramethylcyclopentadienyl t-butylindolyl titanium dichloride, methylenetetramethylcyclopentadienyl di-t-butylindolyl titanium dichloride, methyleneindenyl indolyl titanium dichloride, methyleneindenyl methylindolyl titanium dichloride, methyleneindenyl dimethylindolyl titanium dichloride, methyleneindenyl neindenyl t-butylindolyl titanium dichloride, methyleneindenyl di-t-butylindolyl titanium dichloride, methylenecyclopentadienyl trimethyl phospholyl titanium dichloride, methylenecyclopentadienyl triphenyl phospholyl titanium dichloride, methylencyclopentadienyl phosphoindenyl titanium dichloride, methylenetetramethylcyclopentadienyl trimethyl phospholyl titanium dichloride, methylenetetramethylcyclopentadienyl triphenyl phospholyl titanium dichloride, methylenetetramethylcyclopentadienyl phosphoindenyl titanium dichloride, methyleneindenyl trimethylphospholyl titanium dichloride, methyleneindenyl triphenylphospholyl titanium dichloride, methyleneindenyl phosphoindenyl titanium dichloride and the like.

Other examples are ethylenecyclopentadienyl pyrrolyl titanium dichloride, ethylenecyclopentadienyl methylpyrrolyl titanium dichloride, ethylenecyclopentadienyl dimethylpyrrolyl titanium dichloride, ethylenecyclopentadienyl trimethyl-

pyrrolyl titanium dichloride, ethylenecyclopentadienyl t-butylpyrrolyl titanium dichloride, ethylenecyclope butylpyrrolyl titanium dichloride, ethylenetetramethylcyclopentadienyl pyrrolyl titanium dichloride, ethylenetetramethylcyclopentadienyl methylpyrrolyl titanium dichloride, ethylenetetramethylcyclopentadienyl dimethylpyrrolyl titanium dichloride, ethylenetetramethylcyclopentadienyl trimethylpyrrolyl, titanium dichloride, ethylenetetramethylcyclopenta-5 dienyl t-butylpyrrolyl titanium dichloride, ethylenetetramethylcyclopentadienyl di-t-butylpyrrolyl titanium dichloride, ethyleneindenyl pyrrolyl titanium dichloride, ethyleneindenyl methylpyrrolyl titanium dichloride, ethyleneindenyl dimethylpyrrolyl titanium dichloride, ethyleneindenyl trimethylpyrrolyl titanium dichloride, ethyleneindenyl t-butylpyrrolyl titanium dichloride, ethyleneindenyl di-t-butylpyrrolyl titanium dichloride, ethylenecyclopentadienyl indolyl titanium dichloride, ethylenecyclopentadienyl methylindolyl titanium dichloride, ethylenecyclopentadienyl dimethylindolyl titanium dichloride, ethylenecyclopentadienyl t-butylindolyl titanium dichloride, ethylenecyclopentadienyl di-t-butylindolyl titanium dichloride, ethylenetetramethylcyclopentadienyl indolyl titanium dichloride, ethylenetetramethylcyclopentadienyl methylindolyl titanium dichloride, ethylenetetramethylcyclopentadienyl dimethylindolyl titanium dichloride, ethylenetetramethylcyclopentadienyl t-butylindolyl titanium dichloride, ethylenetetramethylcyclopentadienyl di-t-butylindolyl titanium dichloride, ethyleneindenyl indolyl titanium dichloride, ethyleneindenyl methylindolyl titanium dichloride, ethyl-15 eneindenvi dimethylindolyi titanium dichloride, ethyleneindenvi t-butylindolyi titanium dichloride, ethyleneindenvi di-tbutylindolyl titanium dichloride, ethylenecyclopentadienyl trimethyl phospholyl titanium dichloride, ethylenecyclopentadienyl triphenyl phospholyl titanium dichloride, ethylenecyclopentadienyl phosphoindenyl titanium dichloride, ethylenetetramethylcyclopentadienyl trimethyl phospholyl titanium dichloride, ethylenetetramethylcyclopentadienyl triphenyl phospholyl titanium dichloride, ethylenetetramethylcyclopentadienyl phosphoindenyl titanium dichloride, ethyleneindeand trimethylphospholyl titanium dichloride, ethyleneindenyl triphenylphospholyl titanium dichloride, ethyleneindenyl phosphoindenyl titanium dichloride and the like.

Additional examples are dimethylmethylenecyclopentadienyl pyrrolyl titanium dichloride, dimethylmethylenecyclopentadienyl methylpyrrolyl titanium dichloride, dimethylmethylenecyclopentadienyl dimethylpyrrolyl titanium dichloride, dimethylmethylenecyclopentadienyl trimethylpyrrolyl titanium dichloride, dimethylmethylenecyclopentadienyl t-25 butylpyrrolyl titanium dichloride, dimethylmethylenecyclopentadienyl di-t-butylpyπolyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl pyrrolyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl methylpyrrolyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl dimethylpyrrolyl titanium dichloride, dimethylmethyl enetetramethyl cyclopentadienyl trimethylpyrrolyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl t-butylpyrrolyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl di-t-30 butylpyrrolyl titanium dichloride, dimethylmethyleneindenyl pyrrolyl titanium dichloride, dimethylmethyleneindenyl methylpyrrolyl titanium dichloride, dimethylmethyleneindenyl dimethylpyrrolyl titanium dichloride, dimethylmethyleneindenyl trimethylpyrrolyl titanium dichloride, dimethylmethyleneindenyl t-butylpyrrolyl titanium dichloride, dimethylmethyleneindenyl di-t-butylpyrrolyl titanium dichloride, dimethylmethylenecyclopentadienyl indolyl titanium dichloride, dimethylmethylenecyclopentadienyl methylindolyl titanium dichloride, dimethylmethylenecyclopentadienyl dimethylindolyl 35 titanium dichloride, dimethylmethylenecyclopentadienyl t-butylindolyl titanium dichloride, dimethylmethylenecyclopentadienyl di I-butylindolyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl indolyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl methylindolyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl dimethylindolyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl t-butylindolyl titanium dichloride dimethylmethylenetetramethylcyclopentadienyl di-t-butylindolyl titanium 40 dimethylmethyleneindenyl indolyl titanium dichloride, dimethylmethyleneindenyl methylindolyl titanium dichloride, dimethylmethyleneindenyl dimethylindolyl titanium dichloride, dimethylmethyleneindenyl t-butylindolyl titanium dichloride ride, dimethylmethyleneindenyl di-t-butylindolyl titanium dichloride, dimethylmethylenecyclopentadienyl trimethyl phospholyl titanium dichloride, dimethylmethylenecyclopentadienyl triphenyl phospholyl titanium dichloride, dimethylmethylencyclopentadienyl phosphoindenyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl 45 trimethyl phospholyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl triphenyl phospholyl titanium dichloride, dimethylmethylenetetramethylcyclopentadienyl phosphoindenyl titarium dichloride, dimethylmethyleneindenyl trimethylphospholyl titanium dichloride, dimethylmethyleneindenyl triphenylphospholyl titanium dichloride, dimethylmethyleneindenyl phosphoindenyl titanium dichloride and the like.

Other examples are dimethylsily/cyclopentadieny pyroly titanium dichloride, dimethylsily/cyclopentadieny methylpyroly titanium dichloride, dimethylsily/cyclopentadieny dimethylsily/cyclopentadieny titresthylopyroly titanium dichloride, dimethylsily/cyclopentadieny titresthylsily/cyclopentadieny titresthylsily/cyclopentadieny titresthylsily/cyclopentadieny titresthylsily/cyclopentadieny pyroly titanium dichloride, dimethylsily/tetamethylopyroly titanium dichloride, dimethylsily/tetamethylopyroly titanium dichloride, dimethylsily/tetamethylopyroly titanium dichloride, dimethylsily/tetamethylopyroly titanium dichloride, dimethylsily/tetamethylopyclopentadieny titresthylopyroly titanium dichloride, dimethylsily/tetamethylopyclopentadieny titanium dichloride, dimethylsily/tetamethylopyclopentadieny titanium dichloride, dimethylsily/tetamethylopyclopentadieny titanium dichloride, dimethylsily/tidenyt timethylopyroly titanium dichloride, dimethylsily/tidenyt timethylopyrolytidenyt timethylopyrolytidenytidenytimethylopyrolytidenytidenytimethylopyrolytidenytidenytimethylopyrolytidenytidenytimethylopyrolytidenytidenytimethylopyrolytidenytidenytimethylopyrolytidenytidenytimethylopyrolytidenytidenytimethylopyrolytidenytidenytimethylopyrolytidenytimethylopyrolytidenytidenytimethylopyrolytidenytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytidenytimethylopyrolytimethylopyrolytidenytimethylopyrolytimethylopyrolytimethylopyrolytidenytimethylopyrolytimethylopyrolytimethylopyrolytimethylopyrolytimethylopyrolytimethylopyr

titanium dichloride, dimethylsilykoydopentadienyl dimethylnidolyl titanium dichloride, dimethylsilyhoydopentadienyl tbutylindolyl trainium dichloride, dimethylsilyhopentadienyl di-huylindolyl titanium dichloride, dimethylsilyhteramethyloydopentadienyl rehuylindolyl titanium dichloride, dimethylsilyhteramethyloydopentadienyl trainium dichloride, dimethylsilyhteramethyloydopentadienyl triphenyl phospholyl titanium dichloride, dimethylsilyhteramethyloydopentadienyl t

Additional examples include cyclopentadienyl pyrrolyl zirconium dichloride, cyclopentadienyl methylpyrrolyl zirco-15 nium dichloride, cyclopentadienyl dimethylpyrrolyl zirconium dichloride, cyclopentadienyl tetramethylpyrrolyl zirconium dichloride, cyclopentadienyl t-butylpyrrolyl zirconium dichloride, cyclopentadienyl di-t-butylpyrrolyl zirconium dichloride, pentamethylcyclopentadienyl pyrrolyl zirconium dichloride, pentamethylcyclopentadienyl methylpyrrolyl zirconium dichloride, pentamethylcyclopentadienyl dimethylpyrrolyl zirconium dichloride, pentamethylcyclopentadienyl tetramethvloyrrolyl zirconium dichloride, pentamethylcyclopentadienyl t-butylpyrrolyl zirconium dichloride, pentamethylcyclopentadienyl di-t-butyloyrrolyl zirconium dichloride, indenyl pyrrolyl zirconium dichloride, indenyl methyloyrrolyl zirconium dichloride, indenyl dimethylpyrrolyl zirconium dichloride, indenyl tetramethylpyrrolyl zirconium dichloride, indenyl tbutylpyrrolyl zirconium dichloride, indenyl di-t-butylpyrrolyl zirconium dichloride, cyclopentadienyl indolyl zirconium dichloride, cyclopentadienyl methylindolyl zirconium dichloride, cyclopentadienyl dimethylindolyl zirconium dichloride, cyclopentadienyl t-butylindolyl zirconium dichloride, cyclopentadienyl di-t-butylindolyl zirconium dichloride, pentameth-25 ylcyclopentadienyl indolyl zirconium dichloride, pentamethylcyclopentadienyl methylindolyl zirconium dichloride, pentamethylcyclopentadienyl dimethylindolyl zirconium dichloride, pentamethylcyclopentadienyl t-butylindolyl zirconium dichloride, pentamethylcyclopentadienyl di-t-butylindolyl zirconium dichloride, indenyl indolyl zirconium dichloride, indenyl methylindolyl zirconium dichloride, indenyl dimethylindolyl zirconium dichloride, indenyl t-butylindolyl zirconium dichloride, indenyl di-t-butylindolyl zirconium dichloride, cyclopentadienyl tetramethyl phospholyl zirconium dichloride, cyclopentadienyl tetraphenyl phospholyl zirconium dichloride, cyclopentadienyl phosphoindenyl zirconium dichloride, pentamethylcyclopentadienyl tetramethyl phospholyl zirconium dichloride, pentamethylcyclopentadienyl tetramethyl phospholyl zirconium dichloride, pentamethylcyclopentadienyl phosphoindenyl zirconium dichloride, indenyl tetramethyl phospholyl zirconium dichloride, indenyl tetraphenyl phospholyl zirconium dichloride, indenyl phosphoindenyl zirco-

Additional examples include methylenecyclopentadienyl pyrrolyl zirconium dichloride, methylenecyclopentadienyl methylpyrrolyl zirconium dichloride, methylenecyclopentadienyl dimethylpyrrolyl zirconium dichloride, methylenecyclopentadienyl trimethylpyrrolyl zirconium dichloride, methylenecyclopentadienyl t-butylpyrrolyl zirconium dichloride, methylenecyclopentadienyl di-t-butylpyrrolyl zirconium dichloride, methylenet etramethylcyclopentadienyl pyrrolyl zirconium dichloride, methylenetetramethylcyclopentadienyl methylpyrrolyl zirconium dichloride, methylenetetramethylcyclopentadienyl dimethylpyrrolyl zirconium dichloride, methylenetetramethylcyclopentadienyl trimethylpyrrolyl zirconium dichloride, methylenetetramethylcyclopentadienyl t-butylpyrrolyl zirconium dichloride, methylenetetramethylcyclopentadienyl di-t-butylpyrrolyl zirconium dichloride, methyleneindenyl pyrrolyl zirconium dichloride, methyleneindenyl methylpyrrolyl zirconium dichloride, methyleneindenyl dimethylpyrrolyl zirconium dichloride, methyleneindenyl trimethylpyrrolyl zirconium dichloride, methyleneindenyl t-butylpyrrolyl zirconium dichloride, methyleneindenyl di-tbutylpyrrolyl zirconium dichloride, methylenecyclopentadienyl indolyl zirconium dichloride, methylenecyclopentadienyl methylindolyl zirconium dichloride, methylenecyclopentadienyl dimethylindolyl zirconium dichloride, methylenecyclopentadienyl t-butylindolyl zirconium dichloride, methylenecyclopentadienyl di-t-butylindolyl zirconium dichloride, methylenetetramethylcyclopentadienyl indolyl zirconium dichloride, methylenetetramethylcyclopentadienyl methylindolyl zirconium dichloride, methylenetetramethylcyclopentadienyl dimethylindolyl zirconium dichloride, methylenetetramethylcyclopentadienyl t-butylindolyl zirconium dichloride, methylenetetramethylcyclopentadienyl di-t-butylindolyl zirconium dichloride, methyleneindenyl indolyl zirconium dichloride, methyleneindenyl methylindolyl zirconium dichloride. ride, methyleneindenyl dimethylindolyl zirconium dichloride, methyleneindenyl t-butylindolyl zirconium dichloride, methyleneindenyl dichloride, methy yleneindenyl di-t-butylindolyl zirconium dichloride, methylenecyclopentadienyl trimethyl phospholyl zirconium dichloride, methylenecyclopentadienyl triphenyl phospholyl zirconium dichloride, methylenecyclopentadienyl zirconium dic oindenyl zirconium dichloride, methylenetetramethyloyclopentadienyl trimethyl phospholyl zirconium dichloride, methylenetetramethylcyclopentadienyl triphenyl phospholyl zirconium dichloride, methylenetetramethylcyclopentadienyl phosphoindenyl zirconium dichloride, methyleneindenyl trimethylphospholyl zirconium dichloride, methyleneindenyl triphenylphospholyl zirconium dichloride, methyleneindenyl phosphoindenyl zirconium dichloride and the like.

Other examples include ethylenecyclopentadienyl pyrrolyl zirconium dichloride, ethylenecyclopentasienyl methylpyrrolyl zirconium dichloride, ethylenecyclopentadienyl dimethylpyrrolyl zirconium dichloride, ethylenecyclopentadienyl trimethylpyrrolyl zirconium dichloride, ethylenecyclopentadienyl t-butylpyrrolyl zirconium dichloride, ethylenecyclopentadienyl addienyl addienyl addienyl a tadienyl di-t-butylpyrrolyl zirconium dichloride, ethylenetetramethylcyclopentadienyl pyrrolyl zirconium dichloride, ethyl-5 enetetramethylcyclopentadienyl methylpyrrolyl zirconium dichloride, ethylenetetramethylcyclopentadienyl dimethylpyrrolyl zirconium dichloride, ethylenetetramethylcyclopentadienyl trimethylpyrrolyl zirconium dichloride, ethylenetetramethylcyclopentadienyl t-butylpyrrolyl zirconium dichloride, ethylenetetramethylcyclopentadienyl di-t-butylpyrrolyl zirconium dichloride, ethyleneindenyl pyrrolyl zirconium dichloride, ethyleneindenyl methylpyrrolyl zirconium dichloride, ethyleneindenyl dimethylpyrrolyl zirconium dichloride, ethyleneindenyl trimethylpyrrolyl zirconium dichloride, 10 ethyleneindenyl t-butylpyrrolyl zirconium dichloride, ethyleneindenyl di-t-butylpyrrolyl zirconium dichloride, ethylenecyclopentadienyl indolyl zirconium dichloride, ethylenecyclopentadienyl methylindolyl zirconium dichloride, ethylenecyclopentadienyl dimethylindolyl zirconium dichloride, ethylenecyclopentadienyl t-butylindolyl zirconium dichloride, ethylenecyclopentadienyl di-t-butylindolyl zirconium dichloride, ethylenetetramethylcyclopentadienyl indolyl zirconium dichloride, ethylenetetramethylcyclopentadienyl methylindolyl zirconium dichloride, ethylenetetramethylcyclopenta-15 dienyl dimethylindolyl zirconium dichloride, ethylenetetramethylcyclopentadienyl t-butylindolyl zirconium dichloride, ethylenetetramethylcyclopentadienyl di-t-butylindolyl zirconium dichloride, ethyleneindenyl indolyl zirconium dichloride, ethyleneindenyl methylindolyl zirconium dichloride, ethyleneindenyl dimethylindolyl zirconium dichloride, ethyleneindenyl t-butylindolyl zirconium dichloride, ethyleneindenyl di-t-butylindolyl zirconium dichloride, ethylenecyclopentadienyl trimethyl phospholyl zirconium dichloride, ethylenecyclopentadienyl triphenyl phospholyl zirconium dichloride, ethyl-20 enecyclopentadienyl phosphoindenyl zirconium dichloride, ethylenetetramethylcyclopentadienyl trimethyl phospholyl zirconium dichloride, ethylenetetramethylcyclopentadienyl triphenyl phospholyl zirconium dichloride, ethylenetetramethylcyclopentadienyl phosphoindenyl zirconium dichloride, ethyleneindenyl trimethylphospholyl zirconium dichloride, ethyleneindenyl triphenylphospholyl zirconium dichloride, ethyleneindenyl phosphoindenyl zirconium dichloride and the like.

Additional examples are dimethylmethylenecyclopentadienyl pyrrolyl zirconium dichloride, dimethylmethylenecyclopentadienyl methylpyrrolyl zirconium dichloride, dimethylmethylenecyclopentadienyl dimethylpyrrolyl zirconium dichloride, dimethylmethylenecyclopentadienyl trimethylpyrrolyl zirconium dichloride, dimethylmethylenecyclopentadienyl t-butylpyrrolyl zirconium dichloride, dimethylmethylenecyclopentadienyl di-t-butylpyrrolyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl pyrrolyl zirconium dichloride, dimethylmethylenetetramethyl-30 cyclopentadienyl methylpyrrolyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl dimethylpyrrolyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl trimethylpyrrolyl zirconium dichloride, dimethylmethylmetetramethylcyclopentadienyl t-butylpyrrolyl zirconium dichloride, dimethylmethylmetetramethylcyclopentadienyl di-t-butylpyrrolyl zirconium dichloride, dimethylmethyleneindenyl pyrrolyl zirconium dichloride, dimethylmethyleneindenyl methylpyrrolyl zirconium dichlonde, dimethylmethyleneindenyl dimethylpyrrolyl zirconium dichloride, dimethylmethyleneindenyl trimethylpyrrolyl zirconium dichloride, dimethylmethyleneindenyl t-butylpyrrolyl zirconium dichloride, dimethylmethyleneindenyl di-t-butylpyrrolyl zirconium dichloride, dimethylmethylenecyclopentadienyl indolyl zirconium dichloride, dimethylmethylenecyclopentadienyl methylindolyl zirconium dichloride, dimethylmethylenecyclopentadienyl dimethylindolyl zirconium dichloride, dimethylmethylenecyclopentadienyl t-butylindolyl zirconium dichloride, dimethylmethylenecyclopentadienyl di-t-butylindolyl zirconium dichloride, dimethylmethylenetetramethyl-40 cyclopentadienyl indolyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl methylindolyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl dimethylindolyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl t-butylindolyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl di-t-butylindolyl zirconium dichloride, dimethylmethyleneindenyl indolyl zirconium dichloride, dimethylmethyleneindenyl methylindolyl zirconium dichloride, dimethylmethyleneindenyl dimethylindolyl zirconium dichloride, dimethylmethyleneindenyl 45 neindenyl t-butylindolyl zirconium dichloride, dimethylmethyleneindenyl di-t-butylindolyl zirconium dichloride, dimethylmethylenecyclopentadienyl trimethyl phospholyl zirconium dichloride, dimethylmethylenecyclopentadienyl triphenyl phospholyl zirconium dichloride, dimethylmethylencyclopentadienyl phosphoindenyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl trimethyl phospholyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl triphenyl phospholyl zirconium dichloride, dimethylmethylenetetramethylcyclopentadienyl phosphoindenyl zirconium dichloride, dimethylmethyleneindenyl trimethylphospholyl zirconium dichloride, dimethylmethyleneindenyl triphenylphospholyl zirconium dichloride, dimethylmethyleneindenyl phosphoindenyl zirconium dichloride and the like.

Additional examples are dimethylsilylcyclopentademyl pyrroly zirconium dichloride, dimethylsilylcyclopentademyl methylgyrroly zirconium dichloride, dimethylsilylcyclopentademyl methylgyrroly zirconium dichloride, dimethylsilylcyclopentademyl trional pyrrolyl zirconium dichloride, dimethylsilylcyclopentademyl birulylcyclopentademyl zirconium dichloride, dimethylsilylteramethylcyclopentademyl pyrrolyl zirconium dichloride, dimethylsilylteramethylcyclopentademyl methylgyrrolyl zirconium dichloride, dimethylsilylteramethylcyclopentademyl trimethylgyrolyl zirconium dichloride, dimethylsilylteramethylcyclopentademyl trimethylsilylteramethylcyclopentademyl trimethylgyrolyl zirconium dichloride, dimethylsilylteramethylcyclopentademyl zirconium dichloride, dimethylsilylteramethylcyclopentademylteramethylcyclopenta

silytetramethylcyclopentadienyl di-t-butylpyrrolyl zirconium dichloride, dimethylsilylindenyl pyrrolyl zirconium dimethylsilylindenyl methylpyrrolyl zirconium dichloride, dimethylsilylindenyl dimethylpyrrolyl zirconium dichloride, dimethylsilylindenyl trimethylpyrrolyl zirconium dichloride, dimethylsilylindenyl t-butylpyrrolyl zirconium dichloride, dimethylsilylindenyl di-t-butylpyrrolyl zirconium dichloride, dimethylsilylcyclopentadienyl indolyl zirconium dichloride, dimethylsilylcyclopentadienyl methylindolyl zirconium dichloride, dimethylsilylcyclopentadienyl dimethylindolyl zirconium dichloride, dimethylsilylcyclopentadienyl t-butylindolyl zirconium dichloride, dimethylsilylcyclopentadienyl di-tbutylindolyl zirconium dichloride, dimethylsilyltetramethylcyclopentadienyl indolyl zirconium dichloride, dimethylsilyltetramethylsilyltetramethylcyclopentadienyl indolyl zirconium dichloride, dimethylsilyltetramethylsil tramethylcyclopentadienyl methylindolyl zirconium dichlonde, dimethylsilyltetramethylcyclopentadienyl dimethylindolyl zirconium dichloride, dimethylsilyltetramethylcyclopentadienyl t-butylindolyl zirconium dichloride, dimethylsilyltetramethylcyclopentadienyl di-t-butylindolyl zirconium dichloride, dimethylsilylindenyl indolyl zirconium dichloride, dimethylsilylindenyl methylindolyl zirconium dichloride, dimethylsilylindenyl dimethylindolyl zirconium dichloride, dimethylsilylindenyl t-butylindolyl zirconium dichloride, dimethylsilylindenyl di-t-butylindolyl zirconium dichloride, dimethylsilylcyclopentacienyl trimethyl phospholyl zirconium dichloride, dimethylsilylcyclopentacienyl triphenyl phospholyl pholyl zirconium dichloride, dimethylsitylcyclopentadienyl phosphoindenyl zirconium dichloride, dimethylsityltetramethvicyclopentadienyl trimethyl phospholyl zirconium dichloride, dimethylsilyltetramethylcyclopentadienyl triphenyl phospholyl zirconium dichloride, dimethylsilyltetramethylcyclopentadienyl phosphoindenyl zirconium dichloride. dimethylsilvlindenyl trimethylphosoholyl zirconium dichloride, dimethylsilvlindenyl triphenylphosoholyl zirconium dichloride. ride, dimethylsilylindenyl phosphoindenyl zirconium dichloride and the like.

# 20 2) The catalyst system for polymerizing an olefin.

As the organoaluminum compound (8) constituting the catalyst system for polymerizing an olefin with the above-mentioned transition metal compound (A) in the present invention, a well-known organoaluminum compound can be used. As an example, the organoaluminum compound (81) indicated by the general formula  $R^0_{b}AlZ_{b,b}$  and the cyclic aluminozane (82a) having a structure indicated by the general formula  $L^0_{b}AlZ_{b,b}$  and the cyclic aluminozane (82b) having a structure indicated by the general formula  $R^0_{b}AlZ_{b,b}AlZ_{b,b}$  and  $L^0_{b}AlZ_{b,b}$  and the cyclic aluminozane (82b) having a structure indicated by the general formula  $L^0_{b}AlZ_{b,b}$  and  $L^0_{b}AlZ_{b$ 

Examples of the organoaluminum compound (B1) indicated by the general formula R<sup>3</sup><sub>0</sub>AlZ<sub>5,0</sub> include a triallyslauminum such as trimethylaliminum, triar-propylalaminum, triar-propylaliminum, triar-propylaliminum, triar-propylaliminum, triar-propylaliminum, triar-propylaliminum chloride, displauminum chloride, such as dimethylaliminum chloride, such as dimethylaliminum chloride, such as dimethylaliminum chloride, displauminum chloride, such as dimethylaliminum chloride, displauminum chloride, displau

Next, examples of M<sup>5</sup> and R<sup>5</sup> in the cyclic aluminozane (B2a) having a structure indicated by the general formula {-}(H<sup>5</sup>)-O-}<sub>3</sub>, and/or a linear aluminozane (B2b) having a structure indicated by the general formula R<sup>5</sup>(-A(IR<sup>5</sup>)-O-)<sub>3</sub> AIR<sup>5</sup><sub>2</sub> with a maky group such as methyl, ethyl, n-cycpyl, isopropyl, n-budyl, isobudyl, n-pendyl, nependyl or the like, c. and d are integers of 1 or more. An integer of 1 to 40 is preferred. More preferably, R<sup>5</sup> and R<sup>5</sup> are methyl or isobudyl and c and d are 3 to 20. As the above-mentioned aluminozane, methylatuminozane is operically preferred. The above-mentioned aluminozane can be prepared with various methods, including those well known in the art. For example, the aluminozane can be prepared by contacting a solution of trially/aluminum (for example, trimethylatuminum) in a suitable organic solvent (benzene, alignatic hydrocarbon and the like) with water.

As an another method, there is a method preparing it by contacting a trialkylaluminum (for example, trimethylaluminum and the like) with a crystallized hydrate of a metal salt (for example, a hydrate of cupric sulfate).

In the present invention, in addition to the compound (A) and the compound (B), the catalyst system containing the compound (C) forming an ionic complex by reacting with the transition metal compound can be used.

As the compound (C), any compound forming an ionic complex by reacting with a transition metal compound formed by reacting the compound (A) and the compound (B) can be used.

Preferably, it is a Lewis acid which can furn the transition metal compound into a cation and become a non-coordinating anion corresponding to it or a compound represented by the general formula CPT's (wherein C''s is a cationic oxidizing agent by which the transition metal can be oxidized to become a cation and A' is a non-coordinating anion corresponding to it) or a compound represented by the general formula (L-IPT's (wherein L is a neutral Lewis base, (L-IPT's is a Bronsted acid by which the transition metal can be oxidized to be a cation and A' is a non-coordinating anion corresponding to its.

Preferably, the Lewis acid which can turn the transition metal compound into a cation and become a non-coordinating anion corresponding to it, a compound represented by the general formula C+A or a compound represented by the



general formula (L-H)+A- is a case of a boron compound.

More preferably, it is a boron compound forming an ionic complex by reading with a transition metal compound wherein the above-mentioned Lewis acid is represented by the general formula  $\mathbb{R}^2 \setminus \mathbb{R}^2$ , the compound represented by the general formula  $\mathbb{C}^1 \times \mathbb{R}^2$  is represented by the formula  $\mathbb{C}^1 \times \mathbb{R}^2 \setminus \mathbb{R}^2 \setminus \mathbb{R}^2$ , and the compound represented by the general formula  $(\mathbb{C}^1 + \mathbb{N}^2)$  is represented by the formula  $(\mathbb{C}^1 + \mathbb{N}^2) \times \mathbb{R}^2 \setminus \mathbb{R}^2 \setminus$ 

The example of the Lewis acid which can turn the transition metal compound into a cation and become a non-coorordinating arion responded to it includes tris(pentalfluorophenyl)borane, tris(2,3,5- tentalfluorophenyl)borane, tris(2,3,4-5-tentalfluorophenyl)borane, pris(2,3,4-trifluorophenyl)borane, phenylbis(pentalfluorophenyl)borane and the like.

Tris(pentafluorophenyl)borane is preferred.

Besides, as the example of the compound represented by the general formula C\*(BO,QoQ,Q)\*, the C\* being a cationic oxidizing agent includes a terrosonium cation, an allyf-substituded feroreonium cation, a silvi-substituded feroreonium tetraksic 4.5-triflucorphenyliporate, tetraksic 3.4-triflucorphenyliporate, tetraksic 3.4-triflucorphenyliporate, silvi-substituded feroreonium tetraksic partialful prophenyliporate, silvi-substituded feroreonium tetraksic-partialful prophenyliporate feroreonium tetraksic-partialful prophenylipo

As the example of the compound represented by the general formula (L-H)\*(EO,Q-Q-Q-),\* (L-H)\* being a Bronsted acid includes trialitylammonium, N.N-dialitylaminium, dialitylammonium, triarylphosphonium and the Ilike and the 25 (EO,Q-Q-Q-) being a non-coordinating anion includes the same described above. The example of these combinations includes triethylammonium tetrakis(pentafluorophenyl)borate, tri(h-subylammonium tetrakis(pentafluorophenyl)borate, bir(h-subylammonium tetrakis(pentafluorophenyl)borate, hiv-diventylaminium tetrakis(pentafluorophenyl)borate, N.N-dientylaminium tetrakis(pentafluorophenyl)borate, N.N-dientylaminium tetrakis(pentafluorophenyl)borate, N.N-dientylaminium tetrakis(pentafluorophenyl)borate, N.N-dientylaminium tetrakis(pentafluorophenyl)borate, N.N-dientylaminium tetrakis(pentafluorophenyl)borate, dicyclohexylammonium tetrakis(pentafluorophenyl)borate, bir(methylphenyl)borate, n. dicyclohexylammonium tetrakis(pentafluorophenyl)borate, tridinentylaminium tetrakis(pentafluorophenyl)borate and the like. Ti(rh-butylammonium tetrakis(pentafluorophenyl)borate or N.N-dimetrylaminium tetrakis(pentafluorophenyl)borate or preferred.

### 3) The process for producing an olefin polymer

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In the present invention, as the polymerization catalyst, (1) the above-mentioned compound (A) and the compound (B) or the compound (A), the compound (B) and the compound (C) may be ted into a polymerization vessel with an arbitrary order, or (2) the reacted matter obtained by contacting the above-mentioned compound (A) and the compound (B) and the compound (a) the compound (A), the compound (B) and the compound (b) in advance may be used.

It is desirable to use each component in order that concerning the amount used of each catalyst component in the method mentioned above, the compound (A) may be 0.0001-10 mmol/lip er the capacity of the polymerization vessel, and preferably 0.001-1 mmol/lip the compound (g) at Al atom conversion may be 0.01 to 1000 mmol/lip and preferable 0.101-1000 mmol/lip the compound (C) may be 0.0001 to 20 mmol/l and preferable 0.001 to 20mmol/l, the molar ratio of the compound (B)/the compound (A) may be 0.1 to 10000 and preferably 0.5 to 2000 , and the molar ratio of the compound (C)-the compound (A) may be 0.0 to 1000 and preferably 0.5 to 2000 , and the molar ratio of the compound (C)-the compound (A) may be 0.0 to 1000 and preferably 0.5 to 2000 .

In the present invention, the monomer constituting the oldrin polymer is one or more of oldrins, and an α -oldrins preferably used. The α-oldrin is the one having 2-10 carbon atoms and the example includes eithyrine, propylene, so butner-1, 4-methyl pentene-1, hexane-1, octene-1, winyl cyclohexane and the like. However, the present invention should not be limited to the above-mentioned compound.

The polymerization method should not be limited. For example, a solvent polymerization or slurry polymerization using an aliphatic hydrocarbon such as butane, pentane, hexane, heptane, octane and the like, an aromatic hydrocarbon such as benzene, bulene and the like, or a haliogenated hydrocarbon such as methylenechloride and the like as as solvent (liquid medium), a vapor phase polymerization in the gaseous monomer or the like is can be adopted and either of batchvise polymerization and continuous polymerization are be adopted.

A polymerization temperature can select the range of -50°C to 250°C, but, in particular, the range of -20°C to 100°C is preferred, and a polymerization pressure is preferably within the range of atmospheric pressure to 60 kg/cm². A



polymerization time is generally determined appropriately considering the kind of the polymer aimes apparatus, and a range of 5 minutes to 20 hours can be taken.

In the present invention, a chain transfer agent such as hydrogen and the like in order to control a molecular weight of the polymer can be added.

## EXAMPLE

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The present invention is illustrated in detail according to Examples and Comparative Examples as follows, but the present invention is not limited thereto. Properties of the polymer in Examples were measured by the following methods.

α-olefin content was determined with the characteristic absorption of ethylene and α-olefin by using infrared spectrophotometer (IR-810 manufactured by Nippon Burkou Industry Ltd.) and was represented as the number of short branched-chains per 1000 carbon atterns/SCBI.

A molecular weight and a molecular weight distribution were determined by the following condition with gel permeation chromatograph (150, C manufactured by Waters Company Ltd.).

Column: TSK gel GMH-HT

Measurement temperature was settled at 145°C

Measurement concentration: 10mg/10ml-ODCB

An intrinsic viscosity [n] was measured with Ubbelohde type viscometer at 130°C in tetralin solution. The bigger is the value of intrinsic viscosity [n], the molecular weight of the olefin polymer is the bigger.

The structure of a complex and a ligand was confirmed with <sup>1</sup>H-NMR measurement (R-1500 manufactured by Hitachi Ltd.) and elemental analysis (CHN-O rapid type manufactured by Elemental Company Ltd.).

A measurement apparatus of a melting point (MP-S3 manufactured by Yanaco Company Ltd.) was used for determining a melting point.

#### 25 Example 1

(1) Synthesis of cyclopetadienyl tetramethylpyrrolyl zirconium dichloride

#### (1-1) Synthesis of 2.3.4.5-tetramethylpyrrole

After a 1 liter of four-necked flask equipped with a stirrer, a dropping furnel and a thermometer was substituted with argon, 31g(270mmd) of 3-methyf-2,4-pertanedone, 32g(490mmd) of zinc powder, 150ml of glacial acetic acid was added to 25g(250mmd) of 2.3-butnedione-2-coxime, was transferred into the dropping furnel, and white regulating the temperature of flask at 60°C - 36°C, 230 ml of purified water was added after heading this stury under refluxing for 1 hour. Steam was blown into this mixed solution, and steam distillation was performed. A white solid obtained was filtered and subsequently, it was washed 2 times by using 100 ml of purified water and successively. 2 times by using 100 ml of purified water and successively. 2 times by using 100 ml of purified water and successively. 2 times by using 100 ml of purified water and successively. 2 times by using 10 ml of heading 10 ml of purified water and successively. 2 times by using 10 ml of heading 10 ml of

The melting point of this white solid was 115-11 8°C.

(1-2) Synthesis of 2,3,4,5-tetramethylpyrrolyl lithium salt

After a 100ml of four-necked flask equipped with a stirter, a dropping funnel and a thermometer was substituted with argon, 0.86g (5.4mmol) of 2.3,4.5-tetramethylpyrrole, 30ml of diethyl either were fed and this solution was kept at 5°C. 3.4m(5.4mmol) of n-buryliffitum(1.6 molt) diluted with hexane was dropped into the solution in the flask from the dropping funnel and was kept to be streed for 1 hour at 5°C and for over right at room temperature. The slurry obtained was dried under vacuum and write solid of 2.3,4.5-tetramethylpyrroly lithium saft was obtained.

#### 50 (1-3) Synthesis of cyclopentadienyl tetramethylpyrrolyl zirconium dichloride

After a 100ml of four-necked flask equipped with a stirrer, a dropping funnel and a thermometer was substituted with argon, 1.3g (5.0mmol) of cyclopentacleryl zirconium trichloride and 20ml of toluene were fed and this was kept at -70°C. This sturry was added slowly over a cannula to a sturry wherein 30ml of toluene was added to the above-mensor tioned tetramethylpyrrolyl lithium salt(5.4mmol) to be kept at -70°C. This sturry was kept to be stirred and a temperature was elevated from -70°C to nown temperature for over night. After elevating a temperature, a laquid was eliminated under vacuum and a solid was obtained. 30ml of pentane was added to this solid and a component soluble in pentane was extracted. By fittering the insoluble component, it was separated. By cooling the component soluble in pentane at -20°C, 120mg of a light-yellor cubic orystol or Cyclopentaclienyl treatmenthylpyronyl zirconium dichloride was obtained.



Yield was 6.9%

- <sup>1</sup>H-NMR (C<sub>6</sub>D<sub>6</sub>) data of this cubic crystall was shown as follows.
- δ 6.214 (s, SH), 1.995(s, 6H), 1.819(s, 6H) The result of a simultaneous elemental analysis of C,H,N of this cubic crystal was shown as follows.
- C: 44.3%, H: 5.0%, N: 4.0%
- The melting point of this cubic crystal was 119-135°C.
- (2) Copolymerization of ethylene with α-olefin
- After an autoclave having an inner volume of 400 ml with a sirrer was dried under vacuum and substituted with argon, 170 ml of toluene as a solvent and 30ml of hexner-1 as α-defin were fed and a temperature of the reactor was elevated to 60°C. After temperature up, while regulating ethylene pressure at 6kg/m², ethylene was ted and 50 mmol of methylatuminoxane (MMAC) type 3A manufactured by Toso-Akzo Company Ltd.) was cast in after the inner system became stable. Successively, a solution wherein 50 μml of cytogloratelizing fermethylyproly stronoum dichorida synthesized with the above-mentioned method was dissolved in 5ml of toluene was cast in. Polymerization was carried out for 60 minutes while regulating a temperature at 60°C. As a result of polymerization, ethylene-hexene-1 copolymer having a comonomer content/5080 of 14.6, an intrinsic viscosity [fig) of 2.56 (figl), a molecular weight/Mw) of 179000 and a molecular weight distribution(MwMn) of 2.2 was prepared with a yield of 1.8×10°g per 1 mol of zirconium and 1 hour.

### Example 2

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- Cocolymerization of ethylene with α-olefin
- After an autoclave having an inner volume of 400 m with a stirrer was dried under vacuum and substituted with argon, 170 m for tolume as a solvent and 30m of heamer: as a celefix were fed and a temperature of the reactor was elevated to 80°C. After temperature up, while regulating ethylene pressure at 6kg/cm², ethylene was fed and 1.0 mmol of triisobulytialuminum was cast in after the inner system became stable. Successively, a solution wherein 5.0 jumol of cyclopetatedry tetramethylprofix) stronium dichloride synthesized with the method illustrated in Example 1 (1) was 3 dissolved in 5mil of 10 tiluene was cast into it and subsequently, 15 jumol of tripheny/methyl tetrahids/periat/lucorpheny/l borate was cast in. Polymerization was carried out for 60 minutes while regulating a temperature at 60°C. As a result of polymerization, ethylene-haxene-1 copolymer having 508 of 13.0, [n] of 3.6 (du/g), Mw of 251000 and Mw/Mn of 2.8 was prepared with a yeld of 1.1 x10°D pp or 1 mol of zirconium and in hour.

### 35 Example 3

- (1) Copolymerization of ethylene with α-olefin
- After an autoclave having an inner volume of 400 ml with a silivre was dried under vaccum and substituted with a rgon, 170 ml of toluene as a solvent and 30ml of hexene-1 as c-otelin were fed and a temperature of the reactor was elevated to 60°C. After temperature up, while regulating ethylene pressure at 66µcm², ethylene was fed and 1.0mmol of trisobutylatuminum was cast in eiter the inner system became stable. Successively, a solution wherein 5.0 µmol of cyclopetacing tetramethylyrotyl striconium dichloride synthesized with the method flustrated in Example 1 (1) was solved in 5ml of toluene was cast in and 15 µmol of triphenyimethyl tetrakis(pertatilluorophenyi)borate was cast in after 10 minutes. Polymerization was caried out for 60 minutes while resolution a temperature at 60°C.
  - As a result of polymerization, ethylene-hexene-1 copolymer having SCB of 24.9 and [n] of 3.17(dl/g) was produced with a yield of  $1.4 \times 10^6$  a per 1 mol of zirconium and 1 hour.

#### Example 4

- (1) Copolymerization of ethylene with α-olefin
- After an autoclave having an inner volume of 400 ml with a stirrer was dried under vacuum and substituted with argon, 170 ml of tolusene as a solvent and 30ml of besene-1 as a celefin were fed and a temperature of the reactor was selevated to 60°C. After temperature up, while regulating ethylene pressure at 6kg/cm², ethylene was fed and 1.0mmol of thisobutylaluminum was cast in after the inner system became stable. Successively, a solution wherein 5.9 µmol of cyclopetadiny tetramethylyproly zirconium dichloride synthesized with the method fillustrated in Example 1 (1) was dissolved in 5ml of toluene was cast in and 15 µmol of triphenylmethyl tetraks[pentafluorophenyllborate was cast into it after 30 minutes. Polymerization was performed for 60 minutes while regulating a temperature at 60 km.



As a result of polymerization, ethylene-hexene-1 copolymer having SCB of 25.4 and a high molecular weight insoluble even in tetralin at 130°C was produced with a yield of 1.3×10<sup>6</sup>g per 1mol of zirconium and 1 hour.

#### Example 5

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Copolymerization of ethylene with α-olefin

After an autoclave having an inner volume of 400 ml with a stirrer was dried under vacuum and substituted with argon, 170 ml of tolume as a solvent and 30m of hesene-1 as a celini were fed and the reactor was elevated to 80°C.

10 After temperature up, while regulating ethylene pressure at 6kg/cm², ethylene was fed and 1.0mmol of triisobutylatumnum was cast in after the inner system became stable. Successively, a solution wherein 5.0 µmol of cyclopetadienyl tetramethylymoly zirconium dichloride synthesized with the method illustrated in Example 1 (1) was dissolved in 5ml of toluene was cast in and subsequently, 15 µmol of triphenyfmethyl tetrakis(pentat/luorophenyl)borate was cast in. Polymerization was performed for 60 minutes while regulating a temperature at 50°C.

As a result of polymerization, ethylene-hexene-1 copolymer having SCB of 13.1 and [η] of 2.35(dl/g) was prepared with a yield of 2.9×10<sup>6</sup>g per 1mol of zirconium and 1 hour.

## Example 6

20 (1) Synthesis of pentamethylcyclopentadienyl tetramethylpyrrolyl zirconium dichloride

After a 100m of bur-necked flask equipped with a stirre, a dropping tunnel and a thermometer was substituted with argon, 1.1g (3.8 mmg) of pertuamethylocoppentationally zicconium instituted and this sturry was kept at -70°C. Subsequently, this sturry was added slowly over a connula to a sturry wherein 20m of tollusine was added to the tetramethylyprotry if thins stirry as the starty was reported as the same method of Example (1.1-2) to be kept at -70°C. Under stirring of this sturry, a temperature was elevated from -70°C to room temperature for over night. After temperature up, a liquid was eliminated under vacuum and a solid was obtained. 30m of pertanne was added to this solid and a component soluble in pentane was activated. By liftering the insoluble component, it was separated. By cooling the component soluble in pentane was activated. By liftering the insoluble component, it was separated. By cooling the component soluble in pentane was activated. By liftering the insoluble component, it was separated. By cooling the component soluble in pentane was calinated, visit was 17%.

<sup>1</sup>H-NMR (C<sub>6</sub>D<sub>6</sub>) data of this cubic crystall was shown as follows.

δ 2.034 (s. 15H), 1.956(s. 6H), 1.917(s. 6H)

The result of a simultaneous elemental analysis of C,H,N of this cubic crystal was shown as follows. C: 51.4%. H: 6.6%. N: 3.3%

35 The melting point of this cubic crystal was 180 - 187°C.

(2) Copolymerization of ethylene with α-olefin

After an autoclave having an inner volume of 400 ml with a stirrer was dried under vacuum and substituted with argon, 170 ml of tolume as a solvert and 30ml of havaner-1 as c-delin were fed and a temperature of the reactor was elevated to 60°C. After temperature up, while regulating ethylene pressure at 66/pcm<sup>2</sup>, ethylene was fed and 0.6mnol of trisobulyfaltuminum was cast in after the inner system became stable. Successively, a solution wherein 3.0 µmol of pentamethyf-tyclopetaclieny it termenthyfyrynds 'izronium cichicine's symthesized with the method illustrated above was solved in 3ml of toluene was cast in and subsequently, 9.0 µmol of triphenyfmethyf tetraksfepentalfuorophenyfloorate was cast in. Polymerization was performed for 60 minutes with eroquating a temperature at 60°C.

As a result of polymerization, ethylene-hexene-1 copolymer having SCB of 13.4, [η] of 5.67(dl/g), Mw of 425000 and Mw/Mn of 3.5 was produced with a yield of 4.4× 10<sup>6</sup>g per 1mol of zirconium and 1 hour.

# Example 7

Copolymerization of ethylene with α-olefin

After an autoclave having an inner volume of 400 ml with a stirrer was dried under vacuum and substituted with argon, 170 ml of tolume as a solvent and 30 ml of hexen=1 as the α-defin were fed and a temperature of the reactor was elevated to 80°C. After temperature up, while regulating ethylene pressure at 6kg/cm², ethylene was fed and 0.6mmol of triisobutyleluminum was cast into after the inner system became stable. Successively, the solution wherein 3.0 µmol of pentamethylocylopentadienyl tetramethylpyrrolyl zirconium dichloride synthesized with the method illustrated in Example 6 (1) was dissolved in 3ml of toluene was cast in and subsequently, 9.0 µmol of triphenylmethyl ethylenethy



rakis(pentafluorophenyl)borate was cast in. Polymerization was performed for 60 minutes while regulating a temperature at 80°C.

As a result of polymerization, ethylene-hexene-1 copolymer having SCB of 14.1 and  $[\eta]$  of 3.16(dl/g) was produced with a yield of 3.6×10<sup>6</sup>g per 1mol of zirconium and 1 hour.

# Example 8

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(1)Synthesis of ethylene(1-indenyl)(3-indolyl)zirconium dichloride

#### 10 (1-1) Synthesis of indenyllithium salt

After a \$00ml of four-necked flask equipped with a stirrer, a dropping funnel and a thermometer was substituted with argon, 25g(220mmol) of inclene and 200 ml of diethyl ether were led and this sturry was kept at \$7C. Successively, 140ml (220mmol) of n-butyfithium (1.6moll/) folluted with n-hexane was dropped into a solution in the flask for 160 mins of the flask flas

# (1-2) Synthesis of 1-(3-indolyl)-2-(1-indenyl)ethane

After a 300ml of four-necked flask equipped with a stirrer, a dropping funnel and a thermometer was substituted with argon, 1.6g (7.3mmol) of 3-(2-bromethyl)indole and 60ml of tetrahydrothuran were fed and this solution was kept at 5°C. A solution wherein S0ml of tetrahydrothuran was added to 2.8g(23mmol) of the inderylifibrium stat synthesized as described above was dropped slowly into a solution in the flask for 160 minutes from the dropping hunnel, and was kept to be stirred for 30 minutes at 5°C and for over night at room temperature. From the greenish solution, a liquid was removed under vacuum and an oil was obtained. To this oil, 80ml of toluene and 20ml of purified water were added and a toluene layer was taken out by a separating funnel.

it was extracted with 80ml and 50ml of toluene and the toluene layer joined was washed with 50ml of purified water. The 30 solvent of toluene layer obtained was concentrated under vacuum and 1.8g of a cream-colored crystal of 1-(3-indoly)-2-(1-inden)/ethane was obtained by cooline at 2.0°C.

<sup>1</sup>H-NMR ( $C_8D_8$ ) data of this crystal was shown as follows.  $\delta$  7.698 (m, 2H), 7.200(m), 6.487(s, 1H), 6.077(s, 1H), 3.098(s, 4H), 2.151(s, 1H).

### 35 (1-3) Synthesis of 1-(3-indolyl)-2-(1-indenyl)ethanelithium salt

After a 500ml of four-necked flask equipped with a stirrer, a dropping funnel and a thermometer was substituted with argon, 2.2g(8.4mmol) of 1-(5-holy))-2-(1-indery))ethane and 250 ml of toluene were fed and this solution was kept at 5°C. Successively, 11 ml (17mmol) of n-butylithium (1.6mmol) diluted with n-hazane was dropped into the solution in the flask for 10 minutes from the dropping funnel. This sturry was kept to be stirred for 1 hour at 5°C and for over night at room temperature. This sturry was filtered, a solid provided was washed with 30 ml of toluene and was dried under vacuur.

2.0g of light yellow powder of 1-(3-indolyl)-2-(1-indenyl) ethanelithium salt was obtained. Yield was 88 %.

### 45 (1-4) Synthesis of ethylene(1-indenyl)(3-indolyl) zirconium dichloride

After a 200mf of four-necked flask equipped with a stime; a dropping funnel and a thermometer was substituted with argon, 1.1g (4.1mmol) of 1-(3-indoly)-2-(1-indeny) elbanetifibum salt and 50mf of tetrahydrofuran was refer and this stury was kept at 7:0°C. Subsequently, a sturry wherein 30mf of tetrahydrofuran was added to 1.5gl.4.8mmol) of a complex of zirconium tetrachloride with tetrahydrofuran and kept at 7:0°C was added slowly over a cannulat to the sturry in the flask. This sturry was kept to be stirred and a temperature was elevated from 7:0°C to room temperature for over night. After temperature up, a liquid was removed under vacuum and a solid was obtained. Somi of toluene was added to this solid and a component soluble in the 9th effect. Somi of of childromethane was added to the solid and a component soluble in dichloromethane was extracted to the filtered. 30mf of childromethane was added to see that the solid of the solid solid

δ 7.932(m), 7.288(m), 6.380(s), 3.802(br), 3.118(br), 2.171(s), 1.389(br).



# (2) Copolymerization of ethylene with α-olefin

Polymerization was performed in like manner as Example 6(2) except changing pentamethyloydopentadientyl tetramethyloynoly zirconium dichindro to ethylenel, 'richenyliG-indoly) zirconium dichindro and changing a polymerization time to 10 minutes. As a result of polymerization, ethylene-haxene-1 copolymer haxing SCB of 22.7, [ii] of 1,55(dol.) War, 66000 and MWM/m 01.8 was conclude with a visid of 24-10(2 nor min of zirconium and 1 hour.

### Example 9

### (1) Copolymerization of ethylene with α-olefin

After an autoclave having an inner volume of 400 ml with a stirrer was dried under vacuum and substituted with argon, 170 ml of toluene as a solvent and 30ml of heamer-1 as c-clefin were fed and a temperature of the reactor was devrated to 60°C. After temperature up, while regulating ethylene pressure at 6kg/cm², ethylene was fed and 0.3mmol of trisbothylaluminum was cast in after the inner system became stable. Successively, a solution wherein 1.0 µmol of ethylene (1-indeny)(3-indely) zirconium clirichride symthesized with the method illustrated in Example 8(1) was dissolved in 2ml of toluene was cast in and subsequently, 3.0 µmol of tripheny/methyl tetraksigentat/lucropheny/borate was cast in Chylmerization was performed for of minutes while regulating a temperature at 60°C. As a result of polymerization, ethylene-hazene-1 copolymer having SCB of 12.8 and (n) of 1.84(d/kg) was produced with a yield of 7.7×10.6°g per 1 mol of zirconium and 1 hour.

### Example 10

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### (1) Synthesis of ethylene(1-indenyl)(3-indolyl) titanium dichloride

After a 100ml of four-necked flask equipped with a stirrer, a dropping funnel and a thermometer was substitude mayon, 0.35g (1,3 mmo) of 1-(3-field)? 2-(1-indeny) ethanelithium sail and 20ml of totuene were fed and this slumy was kept at -70°C. Subsequently, a slumy wherein 20ml of totuene was added 10.47g(1.4mmo) of a complex of than num tetrachloride with tetrahydroturan and kept at -70°C was added slowly over a cannula to the slumy in the flask. This slumy was kept to be stirred and a temperature was elevated from -70°C to room temperature for over injith. After temperature up, a liquid was removed under vacuum and a sold was obtained. 30ml of dichloromethane was added to this sold and a component soluble in dichloromethane as edvanted and an insoluble part was separated by fifteing. By cooling the component soluble in dichloromethane at -20°C, 28mg of a black solid of ethylene(1-indeny)(3-indoly)) filanium dichloride was Ostained. Weld was 5.7%.

55 <sup>1</sup>H-NMR (C<sub>5</sub>D<sub>6</sub>) data of this black crystal was shown as follows. δ 7.600(m), 6.438(m), 5.882(br), 3.889(br), 2.942(br), 2.161(s), 1.458(br).

### (2) Copolymerization of ethylene with α-olefin

Polymerization was performed in like manner as Example 6(2) except changing pentamethylcyclopentadienyl tetramethylpyrrolyl zirconium dichloride to ethylene(1-indenyl)(3-indolyl) tetrahum dichloride and changing a polymerization time to 60 minutes. As a result of polymerization, ethylene-hexene-1 copolymer having SCB of 22.9, [h] of 2.45(dic), Mw of 84000 and Mw/Mn of 2.6 was produced with a yield of 1.7×10<sup>10</sup> per 1 mol of tilanium and 1 hour.

### 45 Example 11

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### Copolymerization of ethylene with α-olefin

Polymerization was performed in like manner as Example 7 except changing pentamethylcyclopentadienyl tetramethylcyrrolyl zirconium dichloride to ethylene(1-indeny)(3-indolyl) titanium dichloride and changing a polymerization time to 60 minutes. As a result 4.7x10°p per 1 mol of titanium and 1 hour.

### Comparative Example 1

## (1) Copolymerization of ethylene with α-olefin

After an autoclave having an inner volume of 400 ml with a stirrer was dried under vacuum and substituted with argon, 170 ml of toluene as a solvent and 30 ml of hexene-1 as α-olefin were fed and a temperature of the reactor was

elevated to 60°C. After temperature up, while regulating ethylene pressure at 6kg/cm², ethylene was factor 0.5mmol of triscoutylelluminum was cast in after the inner system became stable. Successively, a solution wherein 2.5 pind dicyclopestading-yi zironium diciloritide was dissolved in 3ml of triblene was cast in and subsequently, 7.5 pind of triplenylmethyl tetraks(pentaliturophenyl)borate was cast in. Polymerization was performed for 60 minutes while regulating a temperature at 60°C.

As a result of polymerization, ethylene-hexene-1 copolymer having SCB of 19.9, [n] of 1.96(dl/g), Mw of 136000 and Mw/Mn of 2.1 was prepared with a yield of 1.6x  $10^7$ g per 1mol of zirconium and 1 hour.

Comparative Example 2

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(1) Copolymerization of ethylene with α-olefin

After an autoclave having an inner volume of 400 ml with a stirrer was dried under vacuum and substituted with argon, 195 ml of toluene as a solvent and 5rd of hexanel 1s ac-defin were find and a temperature of the neactor was 19 elevated to 60°C. After temperature up, while regulating ethylene pressure at RigCom<sup>2</sup>, ethylene was fed and 0.25mmol of triisobutyleluminum was cast in after the inner system became stable. Successively, a solution wherein 0.7 µmol of ethylenebisindenyl zirconium dichloride was dissolved in 3ml of toluene was cast in and subsequently, 2.0 µmol of triphenylmethyl tetrakis/pentratiunorphenyl/borate was cast in. Polymerization was performed to 30 minutes while region to the successive of the subsequently and the subsequently of the

Example 12

(1) Synthesis of cyclopentadienyl (2.5-di-tert-butylpyrrolyl) zirconium dichloride

(1-1)Synthesis of 2.2.7.7-tetramethyl-4.5-epoxyoctane-3.6-dione

into a 2 liter of four-necked flask equipped with a stirrer and a thermometer, 30.3g(169mmol) of 1-bromopinacolone and 800 m of defluy lether were lied and this solution was kept at 20°C. Next, 25.8g(460mmol) of potassium hydrodde 30 was added to the solution in the flask and the solution was stirred for 4 hours at a room temperature. Water of 700 ml at 5°C was added to thus reacted matter to divide into an organic layer and apueous layer. Buethy ether of 300ml was added to the aqueous layer and the liquid of aqueous layer was divided into an organic layer and an aqueous layer. The liquid of organic layer was mixed with the previously obtained liquid of organic layer and the solvent was removed from the mixed liquid under a reduced pressure at 80°C. 11.9g (56 mmol) of yellow solid of 2.2.7.7-tetramethyl-4.5-epoxyoc-35 tam-6.36 flore was obtained; fixed was 66 %.

(1-2) Synthesis of 2,2,7,7-tetramethyl-4-octene-3,6-dione

into a 500m of four-necked flask equipped with a stirrer and a retlux condenser, 11.9g (5emmol) of 2.2,77-tebramethyl-4,5-epoxycotane-3,6-dione and 24.5g (148mmol) of potassium iodide were fed and 300ml of glacial acetic acid was added to these thereby Obtaining a solution. After this solution was refluxed to 5 hours, 1500ml of water was acid to the solution. Thus obtained solution was cooled to 5°C, and a precipitated solid was filtered. After the obtained solid was washed two fines with 10ml of water, and was dissolved with 20ml of effation and was cooled to 2°C. After a solid precipitated was filtered, the crystal was dried under a reduced pressure to obtain 2.7g (14 mmole) of yellow plate crystal of 2.2,77-tetramentyl-4-octere, 36-idione. Yeld was 25-idione. Yeld was 25-idione. Yeld was 25-idione.

<sup>1</sup>H-NMR (C<sub>6</sub>D<sub>6</sub>) data of this yellow crystal was shown as follows.

δ 7.630 (s, 2H), 0.970(s,18H).

(1-3) Synthesis of 2,2,7,7-tetramethyloctane-3,6-dione

Into a 200m of bur-necked flask equipped with a stirrer and a thermometer, 4.0g (20mmol) of 2.7.7-tehramethyl4-octene-3,6-dione and 90ml of glacial acetic acid was fed, and this solution was kept at 20°C. 3.2g (48 mmol) of zinc
powder and 18 ml of water were acided to this solution and this mixture was stirred over night at a room temperature to
only a stirred over night at a room temperature to
of 40 ml was acided to the separated aqueous liquid and was stirred, and an organic layer and an aqueous layer. Pentane
liquid and the previously obtained organic liquid and was stirred, and an organic layer was separated. This organic
liquid and the previously obtained organic liquid ware mixed, and this liquid was washed 3 times with 135 ml of water,
washed once with 135 ml of saturated sodium carbonate aqueous solution and then washed once with saturated
sodium chloride aqueous solution, and was dried with 5 g of sodium suffate for one hour. Sodium sulfate was filtered
from the liquid, and a solvent was evaporated from the liquid under a reduced pressure. 3 2g (16 mmol) of a transparent





oil of 2,2,7,7-tetramethyloctane-3,6-dione was obtained. Yield was 79%.

1H-NMR (C<sub>6</sub>D<sub>6</sub>) data of this transparent oil was shown as follows.

5 2,512 (s. 4H), 1,077(s.18H).

## 5 (1-4) Synthesis of 2,5-di-tert-butylpyrrole

Into a 100ml of four-necked flask equipped with a stirrer, a condenser and a thermometer, 0.5g (3.0mmol) of 2.2.77.4etramethylocaten-3.6-cline and 1.4g (18 mmol) of ammonium acetate were fed, and further, 35 ml of glacial acetic acid was added. This solution was refubred for 6 hours. After cooling to a room temperature, 40 ml of pentane 10 and 60 ml of water were added to the solution thereby to divide the solution into an organic payer and an aqueous layer. The liquid organic layer was westhed with 40 ml of water, and drided with sodium sullate at -20° for over right. Sodium sulfate was filtered from the liquid, and a solvent was removed by exporating from the liquid under a reduced pressure. 0.4g (2.2 mmol) of yellow oil of 2.5-di-let-but/privrice was obtained. Yield was 74% or 10 ml. 10

<sup>1</sup>H-NMR (C<sub>6</sub>D<sub>6</sub>) data of this yellow oil was shown as follows. δ 6.057(s, 1H), 6.009 (s, 1H), 1.223(s, 18H).

## (1-5) Synthesis of 2,5-di-tert-butylpyrrolyl lithium salt

After a 200ml of four-necked flask equipped with a stirrer, a dropping funnel and a thermometer was substituted with a group, 3.7g (21mmol) of 2.5d-lert-but/plymrols, 100ml of hexane were fed and this solution was kept at 7°C. 14m(23 mmol) of n-but/plithium1, for solly diluted with hexane was dropped into the solution in the flask from the dropping funnel and was kept to be stirred for 1 hour at 5°C and for over right at room temperature. This reaction mixture was filtered, and the obtained white solid was washed 2 interes with 30 ml of thexane and was chied under a reduced pressure to obtain 3.0 g of white solid of 2.5d-lert-but/ygrorb(lithium sail. The yield was 80 %.

### 25 (1-6) Synthesis of cyclopentadienyl(2,5-di-tert-butylpyrrolyl) zirconium dichloride

After a 200m of bur-necked flask equipped with a stirrer, a dropping funnel and a thermometer was substitude with argon, 0.7g (3.9mmol) of 2.5-di-tert-bur/pyrrolyl tithium salt and 50ml of tolurane were fed to obtain a sturry and this was kept at -70°C. A folluren(30° m) sturry of 1.1g (4.0 mmol) of cyclopentadienyl zirconium trichloride prepared in another flask was added slowly to the sturry previously prepared, at -70°C. This sturry was kept to be stirred and a termerature was slowly elevated form -70°C to from temperature. After elevating a temperature, a liquid was eliminated under a reduced pressure and a solid was obtained. Pertaine of 50ml was added to this solid and a component soluble in pentaine was extracted. By filtering the insoluble component, it was separated. By cooling the component soluble in pentaine to

35 20°C, 160mg of a white crystal of cyclopentadienyl(2,5-di-tert-butylpyrrolyl) zirconium dichloride was obtained. Yield was 10%.

<sup>1</sup>H-NMR (C<sub>6</sub>D<sub>6</sub>) data of this white crystal was shown as follows. δ 6.634(s, 2H), 6.311 (s, 5H), 1.311(s, 18H).

### (2) Polymerization of ethylene with α-olefin

After an autoclave having an inner volume of 400 ml with a stirrer was dried under vacuum and substituted with argon, 120 ml of toluene as a solvent and 80ml of hexene-1 as c-olelin were fed and a temperature of the reactor was elevated to 60°C. After temperature up, while regulating ethylene pressure at 66µcm², ethylene was fed and 0.5mnod 10 misobulyslauminum was cast in after the inner system became stable. Successively, a solution wherein 2.0 µmol of cyclopentadien/(2.5-di-eth-tolypropty) ziconolim dichorides synthesized with the method illustrated above was dissisted in 5ml of toluene was cast in, and subsequently, 8.0 µmol of triphenylmethyl tetraksigentafluorophenyllborate was cast in. Polymerization was performed for 20 minutes with ergolating a temperature at 60°C.

As a result of polymerization, ethylene-hexene-1 copolymer having SCB of 15.4,  $[\eta]$  of 4.03(dl/g) was produced with a yield of  $2.5 \times 10^6 \eta$  per 1mol of zirconium and 1 hour.

As described above in detail, according to the present invention, by using a catalyst system comprising (A) a transition metal compound(A) represented by the general format (1) and no organoaluminum compound(8) or (A), (B) and a compound(C) forming an ionic complex by reacting with a transition metal compound, an olefin polymer having a high molecular weight can be prepared;

#### ss Claims

 A catalyst component for polymerizing an olefin comprising a transition metal compound (A) represented by the general formula (1) having at least one of a cyclopertadienyl group or a substituted cyclopertadienyl group and at least one of a cyclic licand containing a hetero stom and having a delocalized a bond.

$$\begin{pmatrix} R^1 & R^2 & R^$$

wherein M indicates an element of the Third or the Fourth Group or the Lanthanide Series in the Periodic Table, each of R1 and R2 independently represents a hydrogen atom, a halogen atom, a hydrocarbon or halogenated hydrocarbon group having 1 to 20 carbon atoms, wherein R1 and R2 are the same or different and may be substituted with a group containing a nitrogen atom, a phosphorus atom, an oxygen atom, a sulfur atom or a silicon atom. groups R1 together with the carbon atoms to which they are attached, and generally adiacent to each other, may form a ring, generally a 5- or 6- membered ring, groups R2 together with the groups X to which they are attached, and generally adjacent to each other, may form a ring, generally a 5- or 6-membered ring and R1 and R2 may be bonded together; substituent X indicates an element of the 13th Group, 14th Group or 15th Group in the Periodic Table, each substituent X being the same or different and at least one substituent X being an element other than a carbon atom: Y is a hydrogen atom, a halogen atom or a hydrocarbon group having 1 to 20 carbon atoms; I, m and n are integers, such that 1≤1≤3, 1≤m≤3 and 0≤n≤2; a is 0 or 1; and wherein if M is a metal of the Third Group or a Lanthanide Series element, I+m+n=3 and if M is a metal of the Fourth Group, I+m+n=4.

- 30 2. A catalyst component according to claim 1, wherein M is an element of the Fourth Group of the Periodic Table.
  - 3. A catalyst component according to claim 1 or 2, wherein Y is a halogen atom or an alkyl group having 1 to 8 carbon atoms and n is 2.
- 35 4. A catalyst component according to claim 1, 2 or 3 wherein at least one substituent X is a nitrogen atom or a phosphorus atom.
  - 5. A catalyst system for polymerizing an olefin comprising:

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- an a transition metal compound (A) as claimed in any one of claims 1 to 4, and an organoaluminum compound (B) selected from:
  - (B1) organoaluminum compounds of the general formula R3bAlZ3b and (B2) cyclic aluminoxanes (B2a) of the general formula (-Al(R4)-O-), and linear aluminoxanes (B2b) of the general formula

- wherein each of R3, R4 and R5 independently represents a hydrocarbon group having 1 to 8 carbon atoms, all of R4 and all of R5 may be the same or different; Z is a hydrogen and/or a halogen; b is 0 to 3; and c and d represent an integer of 1 or more.
  - 6. A catalyst system according to daim 5, wherein the organoaluminum compound (B1) is a trialkylaluminum compound.

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A catalyst system according to claim 5, wherein the linear aluminoxane (B2b) is methylaluminoxane.



- A catalyst system according to daim 5, 6 or 7 which further comprises a compound (C) which forms an ionic complex by reacting with a transition metal compound is used in addition to the transition metal compound(A) and the cranscularium compound(B).
- 5 9. A catalyst system according to claim 8, wherein compound (C) is a boron compound.

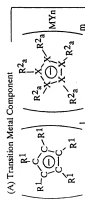
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- A process for producing an defin polymer which process comprises polymerising one or more olefins using a catalvst system according to any one of claims 5 to 9.
- 10 11. A process according to claim 10 wherein the olefin polymer is linear low density polyethylene.

Polymerization Olefin (Co)Polymer



X: Atom of 13th, 14th or 15th Group Y: Hydrogen, Halogen or Alkyl Group

M: Transition Metal of 3rd or 4th Group or Lanthanide Series

(B) Organometallic Component

Organoaluminum Aluminoxane (C) Third Component
Component forming Ionic Complex by
Reacting with Transition Metal Component

Figure 1